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(54) Title: PLANT DEFENSE GENE(S) AND REGULATORY ELEMENT(S)

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(57) Abstract

Novel chitinase gene, and its associated regulatory region, from a monocotyledon plant is described.

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PLANT DEFENSE GENE(S) AND REGULATORY ELEMENT(S)

The present invention relates to regulatory elements functional in plants, especially monocotyledons. In addition, the present invention relates to novel plant genes encoding products involved in plant defense.

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BACKGROUND OF THE INVENTION

The response of plants to microbial attack involves de novo synthesis of an array of proteins designed to restrict the growth of the pathogen. These proteins include hydroxyproline-rich glycoproteins, proteinase inhibitors, enzymes for the synthesis of phytoalexins, enzymes contributing to the reinforcement of cell walls, and certain hydrolytic enzymes such as chitinase and glucanase.

Plant defenses can also be activated by elicitors derived from microbial cell walls and culture fluids. In dicotyledonous plants, extensive studies have shown that 20 microbial attack or elicitor treatment induces the transcription of a battery of genes encoding proteins involved in these defense responses, as part of a massive switch in the overall pattern of gene expression. The functional properties of the promoters of several of these dicotyledonous defense genes have been characterized. In contrast, relatively little is known about the inducible defenses in monocotyledonous plants, including the major cereal crops. For example, the transcriptional regulation of defense genes from monocotyledonous plants has not been examined.

Chitinase (EC 3.2.1.14) catalyzes the hydrolysis of the β -1,4 linkages of the N-acetyl-D-glucosamine polymer chitin. Chitin does not occur in higher plants, but is present in the cell walls of many fungi. Chitinase, which exhibits complex developmental and hormonal regulation, has been found in many species of higher plants. In addition,

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chitinase activity is markedly increased by wounding, ethylene, or microbial elicitors. Furthermore, chitinase is involved in the hypersensitive resistance response to microbial attack. Purified plant chitinase attacks and partially digests isolated cell walls of potentially pathogenic fungi. It is this latter enzyme activity, rather than chitin-binding lectin activity, that is responsible for the inhibition of fungal growth. Chitinase and B-glucanase exhibit synergistic antifungal activity in vitro. A number of pathogenesis-related proteins (also referred to as "PR proteins") have been found to be chitinases or glucanases.

Chitinase genes from a number of dicotyledonous plants (including bean, cucumber, potato, and tobacco) have been isolated and characterized.

Plant chitinases can be divided into at least three classes, based on amino acid sequence and cellular localization. Class I chitinases are basic isoforms which are structurally homologous and are primarily localized in the central vacuole. Basic chitinases contain a catalytic domain, and a cysteine-rich domain similar to rubber hevein. The hevein domain is thought to serve as an oligosaccharide-binding site. There is a variable spacer region between the hevein and the catalytic domains.

Class II chitinases are usually found in the extracellular fluid of leaves and in the culture medium of cell suspensions, suggesting that they are localized in the apoplastic compartment, consistent with a major function in defense. This hypothesis is supported by recent observations that some PR proteins are acidic chitinases.

described cucumber chitinases, show no homology with either Class I or Class II chitinases, but are homologous to a

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lysozyme/chitinase from <u>Parthenocissus quinquifolia</u>. Class III chitinases are located in the extracellular compartment.

While chitinases from dicotyledons have been well characterized, and many of the corresponding genes have been isolated, there is little information available on the structure and expression of chitinase genes from monocotyledons.

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SUMMARY OF THE INVENTION

In accordance with the present invention, we have isolated and characterized a monocotyledon chitinase gene and its associated regulatory sequences. The regulatory sequences of the invention are highly expressed in certain floral organs, and are highly inducible from a low basal level of expression upon exposure to plant defense elicitors.

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The regulatory sequences of the invention are useful, for example, for the controlled expression of a wide variety of gene products, such as reporter constructs, functional proteins (e.g., enzymes), and the like.

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BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 presents a comparison of the amino acid sequences of the invention chitinase (derived from rice)

30 with the amino acid sequences of basic chitinases from dicotyledon plants. The predicted amino acid sequence of RCH10 is shown on the top line, while amino acid sequences of tobacco, potato, and bean basic chitinases are aligned with the RCH10 sequence. Only amino acids differing from the RCH10 sequence are shown. "Dots" indicate gaps in the sequence comparison; while an "*" indicates a stop codon.

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Figure 2 presents a comparison of the amino acid sequence of the RCH10 hevein domain with the amino acid sequences of the hevein domains of other proteins, i.e., rubber hevein [amino acid residues 1-43; see Lucas et al., 5 FEBS Lett. 193: 208-210 (1985)], potato WIN1 and WIN2 [amino acid residues 26-68 of each; see Stanford et al., Genet. 215: 200-208 (1989)], wheat agglutinin isolectin [WGA, amino acid residues 88-127; see Wright et al., Biochemistry 23: 280-287 (1984)], rice RCH10 10 (amino acid residues 22-92), bean basic chitinase [amino acid residues 1-79; see Broglie et al., Proc. Natl. Acad. Sci. USA 83: 6820-6824 (1986)], tobacco basic chitinase (amino acid residues 1-87); tobacco PR-P and PR-Q proteins (amino acid residues 25-57 of each) [see Payne et al., 15 Proc. Natl. Acad. Sci. USA 87: 98-102 (1990) with respect Each of the above to each of the tobacco sequences]. sequences were aligned to maximize sequence identity; only amino acids which differ from the rubber hevein sequence are set forth in the Figure.

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Figure 3 summarizes expression results with RCH10-GUS gene fusions in transgenic tobacco plants. Fig. 3A deals with wound and elicitor induction in leaf tissue; Fig. 3B deals with developmental expression in vegetative 25 organs; and Fig. 3C deals with developmental expression in floral organs.

Figure 4 presents the kinetics of wound and elicitor induction of RCH10-GUS gene fusions in transgenic Fig. 4A presents results using a 30 tobacco leaves. substantially intact promoter (including nucleotides -1512 to +76, with respect to the transcription start site; also presented as nucleotides 374 - 1884 of Sequence ID No. 1, plus nucleotides 1 - 76 of Sequence ID No. 2; referred to 35 as construct BZ4-1); Fig. 4B presents results with a deleted promoter (including only nucleotides -160 to +76, with respect to the transcription start site; also

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presented as nucleotides 1724 - 1884 of Sequence ID No. 1, plus nucleotides 1 - 76 of Sequence ID No. 2; referred to as construct BZ10-1). Open circles designate wounded leaves, while closed circles designate wounded leaves which 5 have also been exposed to elicitor.

Figure 5 summarizes results of RCH10-GUS gene containing 5' deletions to nucleotide (designated as -160 in the Figure, i.e., -160 nucleotide 10 upstream of the translation start site), nucleotide 1810 (designated as -74 in the Figure) and nucleotide 1854 (designated as -30 in the Figure) in transgenic tobacco plants. Panel (A) illustrates wound and elicitor induction of RCH10 promoter deletions in mature leaf tissue. (B) illustrates expression in floral organs. Data are presented mean GUS activities as from replicate determinations with extracts from 3 independent BZ10 (-160) transformants, 14 BZ84 (-74) transformants and 10 BZ10 (-30) transformants.

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DETAILED DESCRIPTION OF THE INVENTION

In accordance with the present invention, there is provided a DNA fragment comprising a monocotyledon promoter characterized as being responsive to physical and/or biological stress; wherein said DNA fragment is further characterized by the following relative pattern of expression in mature plants:

- a low level of expression in leaves;
- 30 a moderate level of expression in plant stems; and

the highest level of expression in the plant roots and in the male and female parts of plant flowers.

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In accordance with another embodiment of the present invention, there are provided DNA construct(s)

comprising the above-described monocotyledon promoter, operatively linked to at least one structural or functional gene, e.g., a reporter gene.

In accordance with yet another embodiment of the present invention, there is provided plant material transformed with the above-described DNA construct(s).

In accordance with still another embodiment of
the present invention, there is provided a method for
inducing the expression of heterologous, functional gene(s)
in monocotyledon and dicotyledon plants, said method
comprising:

subjecting the above-described plant material to 15 conditions which induce transcription of said DNA construct(s).

In accordance with a further embodiment of the present invention, there are provided substantially pure proteins having in the range of about 300 up to 350 amino acids, characterized by:

a hevein domain having in the range of about 40 up to 80 amino acids, wherein said hevein domain is about 70% homologous with respect to dicotyledonous chitinase hevein domains;

a glycine- and arginine-rich spacer region having in the range of about 6 up to 12 amino acids; and

a catalytic domain having in the range of about 240 up to 280 amino acids, wherein said catalytic domain is about 77% homologous with respect to dicotyledenous chitinase catalytic domains.

Proteins of the present invention can optionally further comprise a signal peptide having in the range of about 16 up to 30 amino acids.

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A presently preferred protein of the invention has about 336 amino acids, wherein:

> the hevein domain has about 40 amino acids; the glycine- and arginine-rich spacer region has about 12 amino acids; and

> the catalytic domain has about 262 amino acids.

This presently preferred peptide will optionally have a signal peptide of about 21 amino acids.

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In accordance with a still further embodement of the present invention, there are provided DNA sequences encoding the above-described protein, optionally further containing a readily detectable label.

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In accordance with yet another embodiment of the present invention, there is provided a method for the identification of novel chitinase genes, said method comprising

20 probing a nucleic acid library with at least a portion of the above-described labeled DNA under suitable hybridization conditions, and

> selecting those clones of said library which hybridize with said probe.

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The DNA fragment comprising a monocotyledon contemplated by the promoter present invention responsive to physical and/or biological stress. As used herein, the term "responsive to physical and/or biological 30 stress" refers to DNA sequences which are responsive to exposure to physical stress, such as, for example, wounding (e.g., tearing, folding, bending, and the like), bruising, and the like; or to biological stress, such as, for example, plant defense elicitors (e.g., the high molecular weight fraction heat-released from the cell walls of the soybean fungal pathogen Phytophthira megasperma f. sp. glycinea, purified glucan elicitors, and the like); and so

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forth.

The relative expression pattern of peptides maintained under the expression control of the invention monocotyledon promoter in mature plants is typically as follows:

a low level of expression in leaves;

a moderate level of expression in plant stems; and

the highest level of expression in the plant roots and in the male and female parts of plant flowers.

The monocotyledon promoter of the present invention can be further characterized by reference to the sequences set forth in the Sequence Listing provided herewith, referring specifically to Sequence ID No. 1 (and Sequence ID No. 2). For example, a DNA fragment having substantially the same sequence as nucleotides 1836 to 20 1884, as set forth in Sequence ID No. 1, is operative to confer responsiveness to physical and/or biological stress on a gene associated therewith. Of course, those of skill in the art recognize that longer fragments from the upstream portion of the invention chitinase gene can also 25 be used, such as, for example, a DNA fragment having substantially the same sequence as nucleotides 1810 to about 1884, as set forth in Sequence ID No. 1; a DNA fragment having substantially the same sequence nucleotides 1724 to about 1884, as set forth in Sequence ID 30 No. 1; a DNA fragment having substantially the same sequence as nucleotides 1558 to about 1884, as set forth in Sequence ID No. 1; a DNA fragment having substantially the same sequence as nucleotides 372 to about 1884, as set forth in Sequence ID No. 1; a DNA fragment having 35 substantially the same sequence as nucleotides 1 to about 1884, as set forth in Sequence ID No. 1; and the like.

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addition, sequences downstream of the In transcription start site can also be included in the regulatory elements employed herein (up to about 100 or of nucleotides derived from downstream 5 transcription start site can be employed). Thus, the above-described regulatory elements can be extended to comprise, for example, nucleotides 1 - 76 as set forth in Sequence ID No. 2, thereby forming regulatory constructs such as:

a contiguous sequence of nucleotides comprising nucleotides 1836 to 1884, as set forth in Sequence ID No. 1, plus nucleotides 1 - 76 as set forth in Sequence ID No. 2;

a contiguous sequence of nucleotides comprising nucleotides 1810 to 1884, as set forth in Sequence ID No. 1, plus nucleotides 1 - 76 as set forth in Sequence ID No. 2;

a contiguous sequence of nucleotides comprising nucleotides 1724 to 1884, as set forth in Sequence ID No. 1, plus nucleotides 1 - 76 as set forth in Sequence ID No. 2;

a contiguous sequence of nucleotides comprising nucleotides 1558 to 1884, as set forth in Sequence ID No. 1, plus nucleotides 1 - 76 as set forth in Sequence ID No. 2;

a contiguous sequence of nucleotides comprising nucleotides 372 to 1884, as set forth in Sequence ID No. 1, plus nucleotides 1 - 76 as set forth in Sequence ID No. 2;

a contiguous sequence of nucleotides comprising nucleotides 1 to 1884, as set forth in Sequence ID No. 1, plus nucleotides 1 - 76 as set forth in Sequence ID No. 2;

and the like.

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The monocotyledon promoter of the present invention can be used for the controlled expression (with

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respect to both spatial and temporal expression) of a wide variety of gene products. For example, promoter plus reporter constructs (e.g., wherein said reporter gene is selected from chloramphenical acetyltransferase, β -glucuronidase, β -lactamase, firefly luciferase, and the like) can be used to monitor when and where expression from the invention promoter is induced in a host plant or plant cell.

- the comprising constructs Alternatively, 10 monocotyledon promoter of the present invention, plus structural gene, can be employed for the controlled expression of numerous structural (or functional) genes, such as, for example, the Bacillus thuringensis toxin gene, phytoalexin involved in enzymes encoding 15 genes biosynthesis, proteinase inhibitor genes, lytic enzyme genes, genes encoding inducers of plant disease resistance mechanisms, and the like.
- 20 Plants contemplated for use in the practice of the present invention include both monocotyledons and dicotyledons. Monocotyledons are presently preferred because the invention monocotyledon promoter is expected to be functional in nearly all monocotyledons, whereas dicotyledon promoters have frequently been non-operative when used in monocotyledon hosts. Conversely, it is expected that the invention monocotyledon promoter(s) will be functional in many dicotyledon hosts.
- Exemplary monocotyledons contemplated for use in the practice of the present invention include rice, wheat, maize, sorgham, barley, oat, forage grains, as well as other grains.

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Plants or plant cells containing the above constructs (introduced by standard techniques, such as, for example, by transfection) can be used to study patterns of development, for the controlled expression of various plant defense genes, for the expression of selectable marker genes (to screen for mutants or compounds that modulate stress signal transduction pathways), and the like.

In accordance with one embodiment of the present invention, the rice chitinase structural gene has also been isolated and characterized. This gene is characterized as having only coding sequence (i.e., contains no introns), and encodes the above-described polypeptide, plus signal sequence. The rice chitinase structural gene can be further characterized as having substantially the same nucleic acid sequence as nucleotides +55 through +1062, as set forth in Sequence ID No. 2.

The rice chitinase gene of the present invention 20 encodes a novel protein, i.e., rice basic chitinase. The rice basic chitinase of the present invention can be further characterized as having substantially the same amino acid sequence as amino acids 22 - 357, as set forth in Sequence ID Nos. 2 and 3 (for the mature form of rice basic chitinase) or amino acids 1 - 357, as set forth in Sequence ID Nos. 2 and 3 (for the precursor-form of rice basic chitinase).

Optionally, the rice chitinase structural gene,

or a fragment of at least 100 contiguous nucleotides
thereof, can be labeled (wherein said label is selected
from a radiolabeled molecule, a fluorescent molecule, a
chemiluminescent molecule, an enzyme, a ligand, a toxin, a
selectable marker, etc). The resulting labeled rice

chitinase structural gene (or a portion thereof) can be
used, for example, as a probe (e.g., as part of a method to
identify additional monocotyledon or dicotyledon

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chitinase-like genes), and the like.

One of skill in the art can readily determine suitable hybridization conditions for screening libraries 5 in search of additional monocotyledon or dicotyledon chitinase-like genes. For example, one would preferably use stringent hybridization conditions when screening for other monocotyledon chitinase or chitinase-like genes; while one would likely use milder hybridization conditions 10 when screening for dicotyledon chitinase or chitinase-like Stringent hybridization conditions comprise a temperature of about 42°C, a formamide concentration of about 50%, and a moderate to low salt concentration. mild hybridiation conditions comprise a temperature below 15 42°C, formamide concentrations somewhat below 50%, and Exemplary mild moderate to high salt concentrations. hybridization conditions comprise a temperature of about 37°C, a formamide concentration of about 20%, and a salt concentration of about 5X standard saline citrate (SSC; 20X 20 SSC contains 3M sodium chloride, 0.3M sodium citrate, pH Such conditions will allow the identification of sequences which have a substantial degree of similarity with the probe sequence, without requiring perfect homology for the identification of a stable hybrid. The phrase 25 "substantial similarity" refers to sequences which share at least 50% homology. Preferably, hybridization conditions will be selected which allow the identification of sequences having at least 70% homology with the probe, while discriminating against sequences which have a lower 30 degree of homology with the probe.

In the invention method for inducing gene expression in monocotyledon (and dicotyledon) plants, plant material containing DNA constructs under the expression control of invention monocotyledon regulatory sequences is subjected to conditions which induce transcription of the DNA construct. Such conditions include exposing the plant

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or plant material to physical stress (e.g., wounding) and/or biological stress (e.g., infection, elicitor molecules derived from pathogens).

5 The invention will now be described in greater detail by reference to the following non-limiting examples.

EXAMPLES

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Nucleotide sequences were determined by the dideoxy chain-termination [Sanger et al., PNAS 74: 5463-5467 (1977)]. Fragments for sequencing were obtained by restriction endonuclease digestion or exonuclease III deletion [Ausubel et al., <u>Current Protocols in Molecular Biology</u>, Wiley, NY (1987)].

EXAMPLE I Plant Material

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Rice (Oryza sativa L. cv. IR36) seeds were sterilized in 70% ethanol for 2 minutes and then in a 2% solution of sodium hypochlorite for 30 minutes. Sterilized seeds were germinated and grown in MS medium (without hormones) in darkness [Murashige and Skoog, Physiol. Plant 15: 473-497 25 Two weeks after germination, leaves, roots and stems were harvested separately, then immediately frozen in liquid nitrogen and stored at -80°C until required. (cv. CR76) cell suspension cultures were grown in N6 medium 30 [Chu et al., Scientia Sinica <u>5</u>: 659-668 (1975)] maintained in darkness. The high molecular weight fraction heat-released from mycelial cell walls of Phytophthora megasperma pv. glycinea (Pmg) was used as elicitor [Sharp J. Biol. Chem. <u> 259:</u> 11321-11326 (1984)]. 35 Elicitation experiments were conducted on 5-day-old cultures, the stage of the cell culture cycle during which maximum responsiveness to elicitor was observed.

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EXAMPLE II DNA and RNA Isolation

Genomic DNA from rice cell suspension cultures was prepared according to the method of Ausubel et al., <u>supra.</u> DNA was isolated from tobacco leaves as described by Schmid et al., Plant Cell 2: 619-631 (1990). Plasmid and phage DNA were isolated by standard methods [Maniatis et al., Molecular Cloning: A laboratory manual, Cold Springs Harbor Laboratories, Cold Spring Harbor, NY (1982)]. RNA from cell suspension cultures and plant tissues was prepared by the guanidinium isothiocyanate method [Chomczynski and Sacchi, Anal. Biochem. 162: 156-159 (1989)].

EXAMPLE III Isolation and Characterization of Genomic Rice Clones

A lambda-DASH library containing 15-25 kb genomic fragments from a Sau3A partial digest of rice genomic DNA 20 was a gift from N.H. Chua. pCht12.3, a 650 bp bean basic chitinase cDNA fragment cloned in pBluescript, was used as probe [Hedrick et al., Plant Physiol. 86: 182-186 (1988)]. For library screening, filters were pre-hybridized for 2-4 hours at 42°C in 30% formamide, 5 x Denhardt's solution (1 25 x Denhardt's solution is 0.02% bovine serum albumin, 0.02% Ficoll, and 0.02% polyvinylpyrrolidone), 5 x SSC (1 x SSC is 0.15 M NaCl, 15 mM sodium citrate), and 100 μg of sheared salmon sperm DNA per ml. The filters were then hybridized for 24 hours at 42°C in the same buffer with nick-translated probe DNA. Filters were washed in 2 x SSC, 2% SDS at 42°C for 30 minutes and autoradiographed at -80°C. Purified phage clones containing chitinase sequences were analyzed by restriction endonuclease digestion and Southern blot hybridization. Selected fragments were subcloned into pGEM7 or pBluescript vector.

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EXAMPLE IV DNA Blot Hybridization

Rice genomic DNA samples were digested with various restriction enzymes, fractionated by electrophoresis on a 1% agarose gel and blotted onto a nylon membrane (Genescreen plus). Hybridization to genomic DNA was performed for 24 hours at 65°C in 1% SDS, 1M NaCl, 10% dextran sulfate, 100 µg per ml sheared, denatured salmon sperm DNA, and the DNA probe labeled with [32P]. The membrane was washed with constant agitation, twice in 2 x SSC for 5 minutes at room temperature and once in 2 x SSC, 1% SDS for 45 minutes at 65°C.

Genomic Southern blots with tobacco DNA were probed with the HindIII/SacII fragment of pBI101 containing GUS coding sequences using standard procedures.

EXAMPLE V

20 RNA Blot Hybridization

RNA samples were separated by electrophoresis on a 1% agarose formamide gel in 1 x 3-[N-morpholino]-propanesulfonic acid (MOPS)/EDTA buffer (10 x MOPS/EDTA buffer is 0.5 M MOPS, pH 7.0, 0.01 M EDTA, pH 7.5), and blotted onto a nylon membrane. Before hybridization, the membranes were baked at 80°C for 2 hours. The same hybridization conditions as in Southern blot analysis were used, except that hybridization was at 60° instead of 65°C.

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EXAMPLE VI Fusion Protein Analysis

A 941 bp fragment from the chitinase RCH10 coding region (positions +85 to +1026 relative to the transcription start site; nucleotides 85 - 1026, see Sequence ID No. 2) was inserted into pRX-1, pRX-2, and pRX-

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3 expression vectors [Rimm and Pollard, Gene 75: 323-327 pBZ7-3, generate pBZ7-1, pBZ7-2, and to These plasmids were transferred into respectively. Escherichia coli strain HB101 by the CaCl2 method [Maniatis supra], and the transformed cells grown to stationary phase at 37°C in LB broth. The cells were then inoculated into 5 ml of M9-CA minimal medium containing 100 μ g/ml ampicillin, grown for 3 hours at 37°C, and then induced by addition of indolylacrylic acid to a final 10 concentration of 10 μ g/ml. After 5 hours, the cells were harvested and lysed by sonication in 10 mM TRIS-HCl, pH 8.0, 50 mM EDTA, 8% sucrose, 0.5% Triton X-100, and Soluble bacterial extracts were lysozyme (2 mg/ml). analyzed in a 10% SDS-polyacrylamide gel [Maniatis et al. Immunoblotting was performed as described by 15 <u>supra</u>]. Bradley et al., Planta 173: 149-160 (1988). Antiserum to bean chitinase, prepared employing standard techniques, was obtained as a gift from T. Boller.

20 <u>EXAMPLE VII</u>

Isolation and Nucleotide Sequence of RCH10

A rice genomic library was screened using as a probe the insert of pCht12.3, which contains cDNA sequences of a 25 bean basic chitinase [Hedrick et al., supra]. From 12 clones positive clones, 3 plaque-purified characterized by restriction mapping and Southern blot hybridization. A 2.5 kb HindIII fragment from one of these clones, designated RCH10, was subcloned. 30 sequencing showed that this fragment contained a 1.0 kb open reading frame (ORF), together with 1.5 kb of upstream Subcloning of two HincII fragments that sequence. overlapped the HindIII fragment gave an additional 372 bp of nucleotide sequence 5' of the HindIII fragment and 125 bp 3' of this fragment. This 3.0 kb sequence contained the 35 complete RCH10 chitinase gene (see Sequence ID No. 1).

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single long ORF with no introns encoded polypeptide of 336 amino acids (see Figure 1 and Sequence ID No. 2). Figure 1 shows the primary structure of the RCH10 gene product compared with basic chitinases from 5 dicotyledon plants. The RCH10 polypeptide contains a hydrophobic putative signal peptide of 21 amino acids at the N-terminus, as well as hevein and catalytic domains. The hevein domain of RCH10 is about 40 amino acids long and is cysteine-rich. Figure 2 shows a comparison of the hevein domain of RCH10 with the hevein polypeptide and other gene products containing this domain, including WIN1, WIN2, and wheat germ agglutinin isolectin. The hevein domain of RCH10 shares about 70% amino acid sequence identity with these other hevein domains. 15 domain and catalytic domain of RCH10 are separated by a glycine- and arginine-rich spacer region. The amino acid sequence identity between the RCH10 catalytic domain and the catalytic domains of chitinases from dicotyledons is about 77%.

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EXAMPLE VIII TrpE-RCH10 Fusion Protein

The level of similarity between RCH10 and basic (class I) chitinase genes from dicotyledons strongly suggests that RCH10 encodes a rice chitinase. To confirm the identity of the protein product encoded by the RCH10 gene, a fragment from the coding region (positions +85 to +1026) was inserted into the E. coli expression vectors pRX1, pRX2, 30 and pRX3 to obtain the plasmids pBZ7-1, pBZ7-2, pBZ7-3. pBZ7-1 codes for a fusion polypeptide consisting of 18 amino acids from TrpE, 3 amino acids from the linker sequence, and 314 amino acids from the chitinase gene fused the same reading frame. pBZ7-2 and pBZ7-3 are respectively 1 and 2 bases out of frame compared to pBZ7-1. 35 These three plasmids were transferred into E. coli strain HB101, and soluble bacterial extracts were separated in a

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10% SDS-poly-acrylamide gel and stained with Coomassie blue. The results showed an additional 37.5 kDa polypeptide in the cells transformed with pBZ7-1, whereas no additional polypeptides were detected in cells transformed with pBZ7-2 or pBZ7-3. Western blot analysis showed that the 37.5 kDa species in cells transformed with pBZ7-1 reacted with antiserum to bean chitinase, confirming that the RCH10 gene encodes a rice chitinase.

10 EXAMPLE IX

Transcription Start Site

The transcription start site was determined by primerextension analysis using a synthetic 28-mer oligonucleotide identical to the sequence of the antisense DNA strand at 15 translational the from downstream 132-104 initiation codon (5'-CCG-AAC-TGG-CTG-CAG-AGG-CAG-TTG-G-3'). Primer extension analysis was performed by the method of Jones et al., Cell 48: 79-89 (1987), using the synthetic 20 oligonucleotide wherein the 5' terminus was labeled with [32P]. No band was found in the reaction with RNA isolated from control cells, whereas two bands were detected in the reaction with RNA isolated from elicitor-treated cells. The major product was 186 nucleotides in length and 25 corresponded to the position of the first 'A' in the sequence CCCTCAATCT, which closely resembles an eukaryotic transcription initiator sequence [Smale and Baltimore, Cell 57: 103-113 (1989)]. This position was designated as +1. An additional product two nucleotides smaller than the The putative 30 major reverse transcript was also detected. translational initiation codon was 55 bp downstream from the major transcription start site.

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EXAMPLE X

Flanking Sequences

Putative TATA and CAAT boxes were located 44 and 75 bp 5 respectively upstream from the transcription start site (see Sequence ID No. 1) The DNA sequence between these two boxes was GC-rich (72%). Two inverted putative GC boxes were present at positions -55 to -60 and -66 to -70 [Kadonaga et al., Trends Biochem. Sci. 11: 20-23 (1986)]. 10 A sequence similar to the binding site for an elicitorinducible factor in a parsley phenylalanine ammonia-lyase promoter occurred in the inverted orientation at positions -108 to -117 [Lois et al., EMBO J. 8: 1641-1648 (1989)]. An imperfectly duplicated TGTCCACGT motif was located at 15 positions -752 to -736. In vivo footprinting studies have demonstrated constitutive binding of a nuclear factor to this motif [Lois et al., supra]. Putative cis-acting elements in the 5' flanking region of RCH10 are summarized in Table 1:

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Table 1

5	Repeat seque	nces and putative <u>ci</u> n the RCH10 promoter	<u>s</u> -elements			
	<u>Class</u>	Position*	<u>Sequence</u>			
10	TATA box	1836 - 1843	TATATAA			
	CAT box	1806 - 1810	CCAAT			
15	GC box-like motif	1815 - 1819 1824 - 1830	CGCCC(inverted) CCCGCGG(inverted)			
	Elicitor-inducible PAL footprint	1770 - 1778	TGGCAATGC(inverted)			
20	Constitutive PAL footprint	1133 - 1139 1140 - 1146	TGTCCAA TGTCCAC			
	Direct repeat 1	331 - 343 363 - 374	GTATGTAAAAAG GTATGTAAAAAG			
25	Direct repeat 2	748 - 759 912 - 923	TGGGAGCAGCGG TGGGAGCAGCGG			
30	Direct repeat 3	1459 - 1473 1494 - 1507	TACTCTGTGTGATGA TACT-TGTGTGATGA			
	Inverted repeat 1	541 - 550 1229 - 1238	AATTTTTTAA TTAAAAAATT			
35	Inverted repeat 2	1257 - 1266 1650 - 1659	TCCCCAAGGT TGGAACCCCT			
	Triplicated motif	1723 - 1738	A <u>TGCAT</u> GCATA <u>TGCAT</u>			

Numbers refer to the sequence presented in Sequence ID No. 1

A computer-aided search failed to identify significant sequence homology between the rice RCH10 promoter and the promoter of an ethylene-inducible bean chitinase [Broglie et al., Proc. Natl. Acad. Sci. USA 83: 6820-6824 (1989)]. Two putative polyadenylation signals at positions 1054 (AAATAA; see Sequence ID No. 2) and 1093 (AATAAA; see

^{**} PAL = phenylalanine ammonia-lyase

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Sequence ID No. 2) were found in the 3' flanking region. These sequences fit the consensus polyadenylation sequence (A/GAATAA) described in plants [Heidecker and Messing, Annu. Rev. Plant Physiol. 37: 439-466 (1986)].

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EXAMPLE XI

Organization of Rice Chitinase Genes

To estimate the number of chitinase genes in the rice genome, Southern blots of genomic DNA from rice were hybridized with the SacII-HindIII fragment of pRCH10 (positions 422 to 1021; see Sequence ID No. 2), which encodes a region conserved among class I and class II chitinases. This probe hybridized to several restriction fragments of rice genomic DNA digested with EcoRI, ClaI, HindIII or PvuII, indicating the presence of a family of chitinase genes in the rice genome.

EXAMPLE XII

20 <u>Chitinase Gene Expression in Plants and</u> <u>Elicitor-treated Cell Populations</u>

isolated from rice cell suspension cultures treated with the Pmg fungal elicitor were hybridized with 25 the fragment from the conserved region of the RCH10 gene, and also with an RCH10-specific sequence, the SphI-MluI fragment (positions 114 to 259; see Sequence ID No. 2). A low basal level of chitinase transcripts could be detected in cells of suspension cultures when the fragment from the 30 conserved region was used as probe. However, when the RCH10-specific fragment was used as the probe, no basal level of transcripts was detectable. Thus, the basal level of chitinase transcripts in cells in cultured suspension was not due to RCH10, but represented the expression of 35 other members of the gene family. Following treatment with Pmg elicitor, accumulation of chitinase transcripts could be detected within 2 hours, with maximum levels after 6

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hours. Hybridization with the RCH10-specific probe showed a similar marked accumulation of the RCH10 transcript over the time course of 2-6 hours. Northern blot analysis of RNA from different organs showed that transcripts of rice 5 chitinase accumulate to high levels in roots, but only to barely detectable levels in stems and leaves.

EXAMPLE XIII Construction of Gene Fusions

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A 2538 bp HindIII fragment from the RCH10 gene was subcloned into pGEM7, and a HindIII/Ball fragment (a contiguous fragment containing nucleotides 372 - 1884 of Sequence ID No. 1, plus nucleotides 1 - 76 of Sequence ID No. 2) was then inserted into the HindIII/SmaI site of the GUS expression vector pBI101.2 [Jefferson et al., EMBO J 6: 3901-3907 (1987)] to give pBZ4. A 1463 bp HincII fragment from RCH10 was cloned into the pGEM7 SmaI site, and a XbaI/BalI fragment (a contiguous fragment containing 20 nucleotides 1558 - 1884 of Sequence ID No. 1, plus nucleotides 1 - 76 of Sequence ID No. 2) was then inserted into the XbaI/SmaI site of pBI101.2 to give pBZ14. A 276 bp SphI fragment from RCH10 was cloned into pSP72, and a HindIII/Ball fragment (a contiguous fragment containing 25 nucleotides 1724 - 1884 of Sequence ID No. nucleotides 1 - 76 of Sequence ID No. 2) was then inserted into the HindIII/SmaI site of pBI101.2 to give pBZ10. The RCH10-GUS translational fusions in pBZ4, pBZ14 and pBZ10 were confirmed by direct double-stranded sequencing using 30 a GUS-specific primer.

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EXAMPLE XIV

Tobacco Transformation

pBZ4, pBZ14 and pBZ10 were mobilized from Escherichia

5 coli HB101 into Agrobacterium tumefaciens LBA 4404
[Jefferson et al., supra], and transgenic tobacco plants
generated by the leaf disc method [Rogers et al., Methods
Enzym. 118:627-640 (1986)]. Transformed plants were
selected on Murashige and Skoog medium [Murashige and
O Skoog, supra] containing 200 μg/ml kanamycin and 500 μg/ml
carbenicillin or cefatoxim, and grown at 25°C under a
16-hour light (115 mE)/8-hour dark cycle.

pBZ4 contains the 5' flanking sequence of RCH10 from nucleotide 372 and downstream thereof (i.e., non-coding 15 sequence of 1512 nucleotides), the 55 bp leader sequence and the first 22 bp of the RCH10 coding sequence, fused in frame with the GUS coding sequence in the vector pBl101 [Jefferson et al., supra]. This gene fusion was 20 transferred to tobacco by Agrobacterium tumefaciensmediated leaf disc transformation [Rogers et al., supra] and plants regenerated under kanamycin selection. kanamycin resistant plants, 14 exhibited GUS activity in extracts of young leaves. Twelve of these GUS-positive 25 plants were confirmed as transformants containing one T-DNA copy by Southern blot hybridization, and four, designated BZ4-1, BZ4-5, BZ4-7 and BZ4-14, were selected for further studies.

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EXAMPLE XV

Wound and Elicitor Induction

Discs (about 8 mm in diameter) excised from fully expanded leaves were incubated in 50 mM sodium phosphate buffer (pH 7.0) at 25°C in the dark. Tissue samples were snap frozen in liquid nitrogen and stored at -80°C. Fungal elicitor was the high molecular weight fraction heat-

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released from washed mycelial walls of <u>Phytophthora</u> megasperma f.sp. glycinea [Ayers et al., Plant Physiol. <u>57</u>: 760-765 (1976)], and was applied to wounded tissue in 50 mM sodium phosphate buffer (pH 7.0) at a final concentration of 100 μg glucose equivalents/ml.

Excision wounding of leaf tissue caused a marked increase in GUS activity. In transformants BZ4-1 and BZ4-14, wounding resulted in 10- to 20-fold increases in GUS activity (relative to the low basal levels of 49 and 22 pmole of product/minute/mg protein, respectively, in unwounded tissue; see Figure 3A). In transformants BZ4-5 and BZ4-7, the levels of GUS activity in unwounded leaves were 920 and 570 pmole/minute/mg protein, and wounding caused a 2- to 3-fold increase in these relatively high basal levels.

Addition of fungal elicitor to the leaf tissue immediately after excision caused a further marked stimulation of the expression of the gene fusion, compared with equivalent excision-wounded tissue not treated with elicitor (see Figure 4A). Increased GUS activity was observed 16 hours after elicitor treatment with maximum levels after 48 hours (see Figure 4A), whereas the response to excision wounding in the absence of elicitor was somewhat slower. Overall, elicitor treatment of excised leaf discs caused a 40- to 60-fold increase in GUS activity over low basal levels in BZ4-1 and BZ4-14 plants, compared with a 4- to 6-fold increase in BZ4-5 and BZ4-7 plants, which exhibited higher basal levels of expression (see Figure 3A).

Histochemical analysis of GUS activity <u>in situ</u> showed that wound induction of the gene fusion was restricted to the tissues immediately adjacent to the wound surface, whereas elicitor also induced expression in tissues at a somewhat greater distance from the wound surface.

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Ethylene, administered as ethephon, had no effect on the level of GUS activity in intact leaves.

EXAMPLE XVI

<u>Developmental Expression</u>

In addition to elicitor and wound induction in leaf tissue, the RCH10-GUS gene fusion was also expressed during normal development in the absence of an applied stress. 10 Thus, high levels of GUS were observed in rests and moderate levels in stems compared to the relatively weak expression in young leaves (see Figure 3B). Although there was, as expected, some variation among the independent transformants in the absolute levels of expression, the same overall pattern of GUS activity was observed in each root > stem > leaf. Histochemical analysis showed strong expression of RCH10-GUS in juvenile tissue of apical In stems, GUS staining was localized to the epidermis and vascular system. In the latter, staining was 20 not restricted to specific tissue-types, but was observed in a number of locations including the outer phloem, inner phloem and xylem. No GUS staining was observed in pith or cortical tissue.

25 RCH10-GUS fusion gene also exhibited characteristic pattern of expression in floral organs. Thus while only low levels of GUS activity were observed in sepals and petals, comparable to the levels in leaves from the same plants, relatively high levels were found in 30 anther, stigma and ovary extracts (see Figure 3C). organ-specific pattern of expression was confirmed by histochemical analysis of GUS activity in situ. Moreover, the in situ analysis showed that within anthers there was strong expression of the gene fusion specifically in pollen, since no staining was observed with ruptured 35 anthers from which the pollen had been expelled, whereas strong staining was readily detectable with intact anthers

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containing mature pollen. GUS activity was also directly demonstrated by histochemical staining of isolated pollen.

EXAMPLE XVII

Promoter Deletions

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To localize <u>cis</u>-elements that specify the complex developmental regulation and stress induction of the RCH10 promoter, the expression was analyzed for gene fusions with upstream (i.e., 5') portions of the promoter deleted, e.g., deleted to position 1558 (see Sequence ID No. 1; BZ14); deleted to position 1724 (see Sequence ID No. 1; BZ10); deleted to position 1810 (see Sequence ID No. 1; BZ74); and deleted to position 1854 (see Sequence ID No. 1; BZ30).

Ten independent BZ14 transformants and 7 BZ10 transformants were examined, and in both cases two representative plants were analyzed in further detail.

Strikingly, the full pattern of expression established for the BZ4 plants containing the promoter, deleted only to nucleotide 372 (see Sequence ID No. 1), was also observed in plants containing the much more extensive deletions, i.e., BZ14 (deleted to position 1558, refer to Sequence ID or BZ10 (deleted to position 1724, refer to No. 1) 25 Sequence ID No. 1) See Figure 3B. Thus, the BZ14 and BZ10 transformants exhibited wounding and elicitor induction of GUS activity from low basal levels in leaf tissue, with similar fold-inductions over basal levels and similar absolute levels of GUS activity in induced tissue as 30 observed in BZ4 plants containing the full promoter (containing nucleotides 372 to 1884 as presented Sequence ID No. 1). Likewise, the kinetics for wounding and elicitor induction of the constructs containing substantial promoter deletions (i.e., the 1558 - 1884 and 1724 - 1884 constructs) were the same as with the full The BZ14 and BZ10 plants also showed the same characteristic pattern of expression in floral organs as

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observed with the full promoter, with high levels of GUS activity in anthers, stigmas and ovaries compared to relatively weak expression in sepals and petals (see Figure 3C). In vegetative organs of BZ14 and BZ10 5 transformants, the levels of GUS activity were: root > stem > leaf, as observed with the full promoter, although the expression in roots and stems was markedly reduced compared to BZ4 plants (see Figure 3B).

10 In contrast, deletion of the 5'-most 1724 nucleotides (i.e., to -160 nucleotides from the translation start site) caused a marked reduction in the levels of GUS activity in vegetative organs, although the relative expression in different organs was the same as observed with the full 15 promoter: root > stem > leaf. Thus, there appears to be an enhancer element located between nucleotide 1558 and 1724 that is important for expression in vegetative development, but is not required for floral expression or stress induction.

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To delineate cis-elements in the proximal region of the promoter, floral expression and stress induction of the RCH10-GUS gene fusion were compared in BZ74 and BZ30 transformants. Fourteen BZ74 and 10 BZ30 transformants were examined. BZ74 (i.e., where 5'-noncoding nucleotides from 1810 and upstream thereof are deleted) transformants still exhibited wounding and elicitor induction of GUS activity from low basal levels in leaf tissue, although the absolute induction was not as high as in BZ10 plants However, BZ30 (i.e., where 5'-noncoding (Figure 5A). nucleotides from 1854 and upstream thereof are deleted) transformants showed no increase in GUS activity response to wounding and elicitor treatment, indicating the presence of a cis-element for stress induction between 35 nucleotide 1810 and 1854 (Figure 5A). In contrast, deletion of the first 1810 upstream nucleotides abolished expression in floral organs (Figure 5B), indicating the

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presence of a distinct cis-element necessary for floral expression but not stress induction located between nucleotide 1724 and 1810.

5 <u>EXAMPLE XVIII</u>

GUS Assays

GUS activity was assayed in tissue extracts by production the determination of fluorimetric 10 4-methylumbelliferone from the corresponding B-glucuronide [Jefferson et al. supra; Jefferson, Plant Mol. F. ol. Rep. Root, stem and leaf tissues were <u>5</u>: 387-405 (1987)]. collected from 10 cm-tall plantlets and floral organs were Protein was collected from mature fully open flowers. 15 determined by the method of Bradford [Anal. Biochem. 72: 248-254 (1976) and GUS activity was expressed as pmole of product/minute/mg of protein. Histochemical localization of GUS activity in situ was performed with the chromogenic B-D-glucuronide 5-bromo-4-chloro-3-indolyl substrate Stem sections were cut by hand, vacuum-20 (X-gluc). infiltrated with 50 mM sodium phosphate buffer (pH 7.0) containing X-gluc and incubated at 37°C. Flowers and roots were directly incubated in X-gluc solution. overnight incubation, chlorophyll was removed by immersion 25 of the tissue samples in 70% ethanol prior to examination using a Nikon Diaphot TMD microscope.

While the invention has been described in detail with reference to certain preferred embodiments thereof, it will be understood that modifications and variations are within the spirit and scope of that which is described and claimed.

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SUMMARY OF SEQUENCES

Sequence ID No. 1 is the nucleic acid sequence for a regulatory region (i.e., the upstream or 5'region) of a rice chitinase gene of the invention.

Sequence ID No. 2 is the nucleic acid sequence and deduced amino acid sequence for a rice chitinase gene according to the present invention.

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Sequence ID No. 3 is the deduced amino acid sequence for the rice chitinase gene presented in Sequence ID No. 2.

SEQUENCE LISTING

- (1) GENERAL INFORMATION:
 - (i) APPLICANT: LAMB, Ph.D., CHRISTOPHER J. ZHU, Ph.D., QUN
 - (ii) TITLE OF INVENTION: PLANT DEFENSE GENES AND PLANT DEFENSE REGULATORY ELEMENTS
 - (iii) NUMBER OF SEQUENCES: 3
 - (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: PRETTY, SCHROEDER, BRUEGGEMANN & CLARK
 - (B) STREET: 444 South Flower Street, Suite 2000
 - (C) CITY: Los Angeles
 - (D) STATE: California
 - (E) COUNTRY: United States
 - (F) ZIP: 90071-2921
 - (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PG-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25
 - (vi) PRIORITY APPLICATION DATA:
 - (A) APPLICATION NUMBER: US 07/704,288
 - (B) FILING DATE: 22-MAY-1991
 - (C) CLASSIFICATION:
 - (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: Reiter, Mr., Stephen E.
 - (B) REGISTRATION NUMBER: 31192
 - (C) REFERENCE/DOCKET NUMBER: FP41 9322
 - (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: (619) 546-7437
 - (B) TELEFAX: (619) 546-9392
 - (C) TELEX: 9103330318
 - (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1884 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: unknown
 - (ii) MOLECULE TYPE: DNA (genomic)
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

GTTAACTGCC	AGCTTCAAAT	TATTTATAGA	TAATTTAATA	GCCAATTCA	CTAATAGTTA	60
TTTATTATAC	TATTAATATC	TGATCTCACC	TGAGTCATAC	TACAGCTGG	TACAAATGTG	120
TAGTGTACTA	стстттстст	CTTCTTTAT	CTCTTTAAAA	TATGTTATAC	G CGGCTTATAA	180
CTGTTATTGT	ACCTGCTCTA	AGTCGATCGT	GATGATCGAT	CATTCGTCA	ATGTTACCAC	240
GTCCAGTGAC	TTATCCATGG	TTCACCTTAC	TAAAAAAT	GATTTTTATO	GACAACTCCT	300
TTAATTTTGT	TCAAACGGAC	CAAAGAAACC	CGTATGTAAA	AAGGTTGGGA	ATATCTGATC	360
CTGTATGTAA	AAAGCTTGGA	ATATCTGATA	GAGGGCAAAC	TTGTGAAAAT	TGTTTTTTA	420
AGATGGACCT	CTTAACAAGC	CTACTTGCAA	AAAATCGACC	TATTTACATA	GACGGACTTG	480
TTAAGAGACT	TGTCTATGAA	AATCGGTGGA	TAGCATGACC	GGTCACAATA	CTTCCCCTAT	540
AATTTTTAA	TCCTCCTAGA	TAAACCCTAT	CTCTCTCTTC	ATGTTCTTTG	CTTTCCATCT	600
ATAGTCTCGC	ATCCCTCATC	ACCTCCCATT	CCTCTCTCTC	TCACCCCCTG	CTCAGTGGGA	660
GCGCAGCTGG	CGATGGCACC	ACCGGCGACA	AGAGGGGCCA	GAGGCTAGCA	TGTGCACGGA	720
AGTGACAATG	GCGCCACATG	ATTAGCATGG	GAGCAGCGGC	GCGTTTCATC	AGGACACGCT	780
GCAATTGGCT	CTAGTGACGG	CACCCTTGAG	AGGACATGGT	AGCGGTGGCG	CCTCAGGAGT	840
GGTGGGGCAC	GGTGGCAGAA	CTCCGGCGGT	GGCAAGCCAC	CACACAGCGA	CAGATCCACC	900
ACCACCGACC	TTGGGAGCAG	CGGGGCCTCA	GCGGTGATGA	CGATGGTAGA	TCGAAGCTAG	960
GGTTTCTATT	TTTTTTTGCT	GCAAAAATCA	CTTTTTACAC	ATGGGTACAT	GCATGTTTTT	1020
TACATACACC	TAGTATTAGG	TGGGCCGTCC	ACCCGTTCGC	AAAGATCATT	TATGCAGTCA	1080
TCATGATCGG	AGATGGAACT	ATGGAGACAT	ATATGCAAGT	ATTTGGCCAA	CATGTCCAAT	1140
GTCCACCAGA	TTGGGAGCTC	AATCCTACCC	CGTGGTATGG	GTATGTTACT	GTGCGCCTAA	1200
TATTTACGTA	CGCTGGTTTA	ATCTATTTT	AAAAAATTTG	CTACATACTC	CCTCCGTCCC	1260
CAAGGTTGGC	TTTTTTTTT	TGGAGGGAGA	GAGTAATATT	TAGAGTTTGT	GGTTTTTGTT	1320
ATTGAACACC	TTAAAAGGCA	TGAAACGACT	TGTCGGAGAA	CGAATCTCCT	CTAGCAGGGA	1380
AGCAACGAAC	CTCCCAAAAA	AAACAAAAA	AAACTCCTCC	TTTCATGATT	CAACCAAAGG	1440
GCAATTTGAG	ATCGAGCCTA	CTCTGTGTGA	TGAACTCAAA	ACACAATCAA	GTATACTTGT	1500
STGATGAGCG	GTGAGCCAGA	TATGTTCCTG	CTCTGTCCGT	GCTCGACTCA	ATTCATTGTC	1560
ACCCTAGCG	ATTTCCATTA	ATGCAATGAC	TATATGAAAT	GCAAAGATGT	ACTATATGAC	1620
ACTAGTTGG	ATGCACAATA	GTGCTACTAT	GGAACCCCTT	TTGCCCCTCT	AATAGTAGGA	1680

				• OTT • C	TAAC C	GATGCATG	C AT	ATGC	ATGA	1740
TCTAGGCTA	A ATGACG	TTTC AAT	AAATCAG	AGIIAG		ттсттстт	'G AA	AAAT	ACCA	1800
TATGTGAGT	G TCTGTI	AATC GTG	GCAAATT	GGCAAT	rgcaa i	11611611		TCCC	ATCG	1860
AGATGCCAA	T ACTACO	CCCA CTI	CCCGCGG	CGCTCI	A TATA	AAGCCATC	ic Gu	1000	AIOO	1884
CTTCTTCCT	C ACAAA	TTTC CCI	CC							1004
(2) INFOR	MATION I	FOR SEQ	D NO:2:							
(i)	(A) LEI (B) TYI	E CHARACT NGTH: 11: PE: nucle RANDEDNE: POLOGY: 1	eic acid S: unkno	Jarra						
(ii)	MOLECUL	E TYPE:	DNA (gene	omic)						
(ix)	FEATURE (A) NA (B) LO	: ME/KEY: CATION:	CDS 551062							
		E DESCRI	DTION. S	EO ID	NO:2:					
(xi)	SEQUENC	ATACA GO	CAACTCAGO	GATCT	TATAT	TTACCCA	CA C	ACC	ATG Met 1	57
AGA GCG	CTC GCT	GTG GTG	GCC ATG	GIG GG	CC AGG	000 TTC	стс	ccc		
	Leu Ala 5	Val Val	Ala Met	10			15			105
	Leu Ala 5 CAT GCC His Ala	Val Val GAG CAG Glu Gln	AIA MEC	10	AC CCC	ccc ccc	15 GCG	GTG	TGC	105 153
GCC GTG Ala Val CCC AAC Pro Asn	CAT GCC His Ala 20	Val Val	TGC GGC Cys Gly 25	10 AGC CA	AG GCC In Ala	GGC GGC Gly Gly 30	GCG Ala	GTG Val	TGC Cys TCC	
GCC GTG Ala Val CCC AAC Pro Asn 35 GAC TAC Asp Tyr	CAT GCC His Ala 20 TGC CTC Cys Leu	GAG CAG Glu Gln	TGC GGC Cys Gly 25 AGC CAG Ser Gln 40 TGC CAG Cys Gln	10 AGC CASER G:	AG GCC In Ala	GGC GGC Gly Gly 30 TGC GGC Cys Gly 45	GCG Ala TCC Ser	GTG Val ACC Thr	TGC Cys TCC Ser	153
GCC GTG Ala Val CCC AAC Pro Asn 35 GAC TAC Asp Tyr	CAT GCC His Ala 20 TGC CTC Cys Leu TGC GGC Cys Gly	GAG CAG Glu Gln TGC TGC Cys Cys GCC GGA Ala Gly	TGC GGC Cys Gly 25 AGC CAG Ser Gln 40 TGC CAG Cys Gln	AGC CASET G	AG GCC TGG TTP CAG TGC CON AGC	GGC GGC GGC GGC GGC GGC GGC GGC	GCG Ala TCC Ser CTG Leu	GTG Val ACC Thr CGG Arg	TGC Cys TCC Ser CGG Arg 65	153 201

		Ala				Thr				Val	GCC Ala	393	
	Ala								Ala		AAC Asn	441	
Arg					Phe			Ser			ACC Thr 145	489	
				Ala							TGC Cys	537	
							Pro				AGC Ser	585	
		Pro					AAG Lys					633	
							GGG Gly					681	
							CTC Leu					729	
							TGG Trp 235					777	
							GGC Gly					825	
							GGC Gly					873	
							CAT His					921	
			Gly				TAC Tyr					969	
		Ala				Tyr	AGC Ser 315					1017	

CCT AAG CTT CGC CTA CCT AGC TTC CAC ACA GTG ATA AAT AAT CAC Pro Lys Leu Arg Leu Pro Ser Phe His Thr Val Ile Asn Asn His 325 330 335
TGATGGAGTA TAGTTTACAC CATATCGATG AATAAAACTT GATCCGAATT CTCGCCCTAT
AGTGAGTCGT ATTAGTCGAC AGCTCTAGA
(2) INFORMATION FOR SEQ ID NO:3:
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 336 amino acids(B) TYPE: amino acid(D) TOPOLOGY: linear
(ii) MOLECULE TYPE: protein
(x1) SEQUENCE DESCRIPTION: SEQ ID NO:3:
Met Arg Ala Leu Ala Val Val Ala Met Val Ala Arg Pro Phe Leu Ala 1 5 10 15
Ala Ala Val His Ala Glu Gln Cys Gly Ser Gln Ala Gly Gly Ala Val 20 25 30
Cys Pro Asn Cys Leu Cys Cys Ser Gln Phe Gly Trp Cys Gly Ser Thr 35 40 45
Ser Asp Tyr Cys Gly Ala Gly Cys Gln Ser Gln Cys Ser Arg Leu Arg 50 55 60
Arg Arg Arg Pro Asp Ala Ser Gly Gly Gly Gly Ser Gly Val Ala Ser 65 70 75 80
Ile Val Ser Arg Ser Leu Phe Asp Leu Met Leu Leu His Arg Asn Asp 85 90 95
Ala Ala Cys Pro Ala Ser Asn Phe Tyr Thr Tyr Asp Ala Phe Val Ala 100 105 110
Ala Ala Ser Ala Phe Pro Gly Phe Ala Ala Ala Gly Asp Ala Asp Thr 115 120 125
Asn Lys Arg Glu Val Ala Ala Phe Leu Ala Gln Thr Ser His Glu Thr 130 135 140
Thr Gly Gly Trp Ala Thr Ala Pro Asp Gly Pro Tyr Thr Trp Gly Tyr 145 150 155 160
Cys Phe Lys Glu Glu Asn Gly Gly Ala Gly Pro Asp Tyr Cys Gln Gln 165 170 175
Ser Ala Gln Trp Pro Cys Ala Ala Gly Lys Lys Tyr Tyr Gly Arg Gly 180 185 190

Pro	Ile	Gln 195	Leu	Ser	Tyr	Asn	Phe 200	Asn	Tyr	Gly	Pro	Ala 205	G1y	Gln	Ala
Ile	Gly 210	Ala	Asp	Leu	Leu	Gly 215	Asp	Pro	Asp	Leu	Val 220	Ala	Ser	Asp	Ala
Thr 225	Val	Ser	Phe	Asp	Thr 230	Ala	Phe	Trp	Phe	Trp 235	Met	Thr	Pro	Gln	Se1 240
Pro	Lys	Pro	Ser	Cys 245	Asn	Ala	Val	Ala	Thr 250	Gly	Gln	Trp	Thr	Pro 255	Ser
Ala	Asp	Asp	Gln 260	Arg	Ala	Gly	Arg	Val 265	Pro	G1y	Tyr	Gly	Val 270	Ile	Thr
Asn	Ile	Ile 275	Asn	Gly	Gly	Leu	Glu 280	Cys	Gly	His	Gly	Glu 285	Asp	Αs	Arg
Ile	Ala 290	Asp	Arg	Ile	Gly	Phe 295	Tyr	Lys	Arg	Tyr	Cys 300	Asp	Ile	Leu	Gly
Val 305	Ser	Tyr	Gly	Ala	Asn 310	Leu	Asp	Cys	Tyr	Ser 315	Gln	Arg	Pro	Ser	Ala 320
Pro	Pro	Lys	Leu	Arg 325	Leu	Pro	Ser	Phe	His 330	Thr	Val	Ile	Asn	Asn 335	His

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That which is claimed is:

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- 1. A DNA fragment comprising a monocotyledon promoter characterized as being responsive to physical and/or biological stress; wherein said DNA fragment is further characterized by the following relative pattern of expression in mature plants:
 - a low level of expression in leaves;
 - a moderate level of expression in plant stems; and
- the highest level of expression in the plant roots and in the male and female parts of plant flowers.
- 2. A DNA fragment according to Claim 1 having substantially the same sequence as nucleotides 1836 to about 1884, as set forth in Sequence ID No. 1.
- 3. A DNA fragment according to Claim 2 further comprising, as part of the same contiguous fragment, substantially the same sequence as nucleotides 1 76, as 20 set forth in Sequence ID No. 2.
 - 4. A DNA fragment according to Claim 1 having substantially the same sequence as nucleotides 1810 to about 1884, as set forth in Sequence ID No. 1.

5. A DNA fragment according to Claim 4 further comprising, as part of the same contiguous fragment, substantially the same sequence as nucleotides 1 - 76, as set forth in Sequence ID No. 2.

6. A DNA fragment according to Claim 1 having substantially the same sequence as nucleotides 1724 to about 1884, as set forth in Sequence ID No. 1.

7. A DNA fragment according to Claim 6 further comprising, as part of the same contiguous fragment, substantially the same sequence as nucleotides 1 - 76, as set forth in Sequence ID No. 2.

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- 8. A DNA construct comprising the monocotyledon promoter of Claim 1 operatively linked to at least one reporter gene.
- 9. A DNA construct according to Claim 8 wherein said reporter gene is selected from chloramphenicol acetyltransferase, β -glucuronidase, β -lactamase, or firefly luciferase.
- 10. A DNA construct comprising the monocotyledon promoter of Claim 1 operatively linked to at least one structural gene.
- 11. A DNA construct according to Claim 10 20 wherein said structural gene is selected from the Bacillus thuringensis toxin gene, genes encoding enzymes involved in phytoalexin biosynthesis, proteinase inhibitor genes, lytic enzyme genes, genes encoding fungal elicitors, or genes encoding inducers of plant disease resistance mechanisms.

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- 12. Plant material containing the DNA construct of Claim 8.
- 13. Plant material containing the DNA construct 30 of Claim 10.
 - 14. A method for inducing the expression of heterologous, functional gene(s) in monocotyledon and dicotyledon plants, said method comprising:
- subjecting the plant material of Claim 13 to conditions which induce transcription of said DNA construct.

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- 15. A substantially pure protein having in the range of about 300 up to 350 amino acids, characterized by:
 - a hevein domain having in the range of about 20 40 amino acids, wherein said hevein domain is about 70 % homologous with respect to dicotyledonous chitinase hevein domains;
 - a glycine- and arginine-rich spacer region having in the range of about 6 up to 12 amino acids; and
- a catalytic domain having in the range of about 240 280 amino acids, wherein said catalytic domain is about 77 % homologous with respect to dicotyledenous chitinase catalytic domains.
- 16. A protein according to Claim 15 having substantially the same amino acid sequence as set forth in Sequence ID No. 3.
- 20 17. A DNA encoding a protein according to Claim 15.
 - 18. A DNA according to Claim 17 wherein said DNA further contains a readily detectable label.
 - 19. A DNA according to Claim 18 wherein said label is selected from a radiolabeled molecule, a fluorescent molecule, a chemiluminescent molecule, an enzyme, a ligand, a toxin, or a selectable marker.
 - 20. A method for the identification of novel chitinase genes, said method comprising
- probing a nucleic acid library with at least a portion of the DNA of Claim 18 under hybridization conditions, and
 - selecting those clones of said library which hybridize with said probe.

FIG. |∆

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HEVEIN DOMAIN

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CATALYTIC DOMAIN

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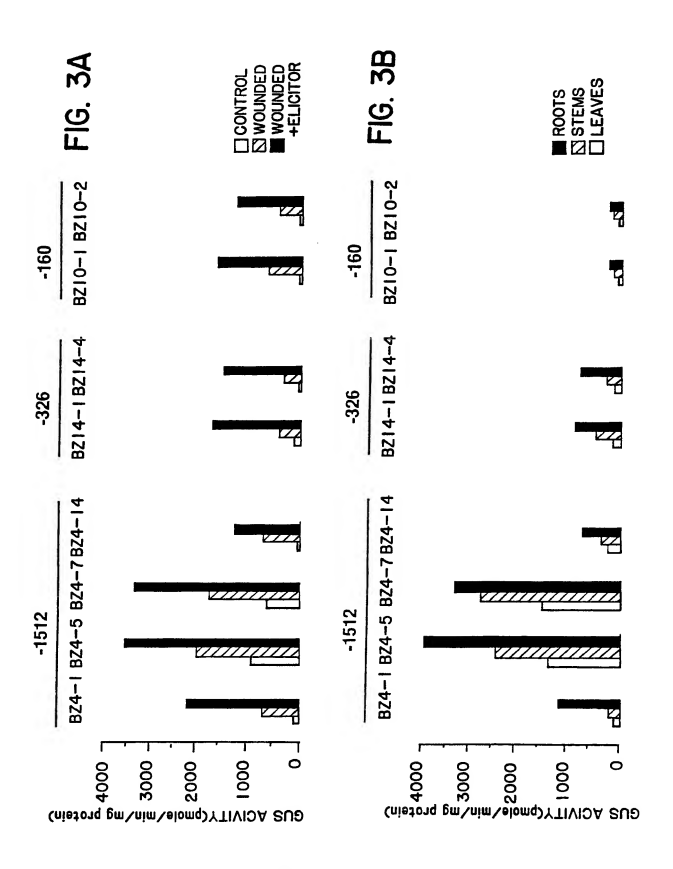
Q -GGPSPAP----TDLSAL I-SRSTFDQMLK Q PGGPTPTPPGGGDLGS I I-SSSMFDQMLK

Q SRIRRRPDASGGGSGVASIVSRSLFDLWLL

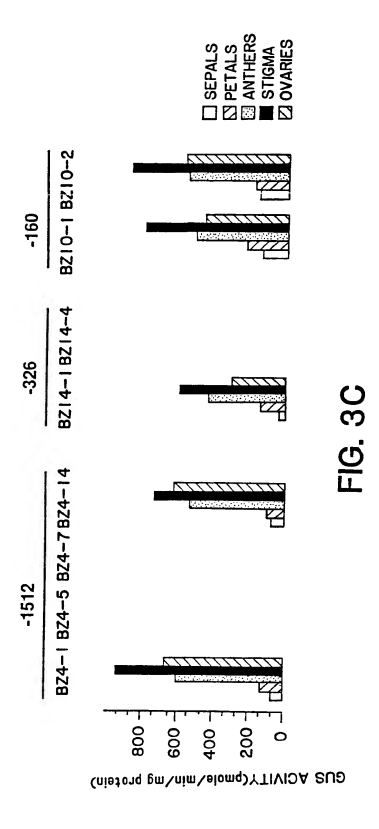
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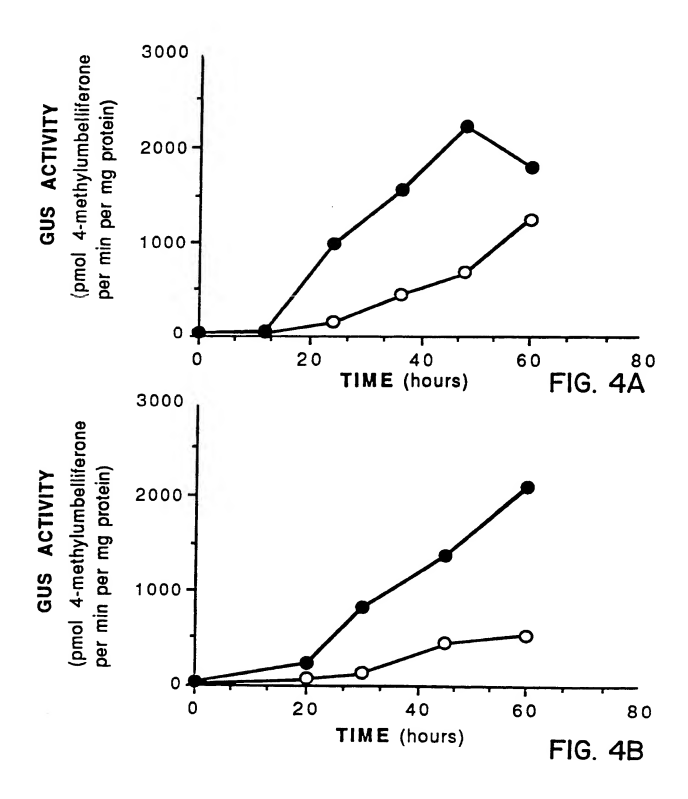


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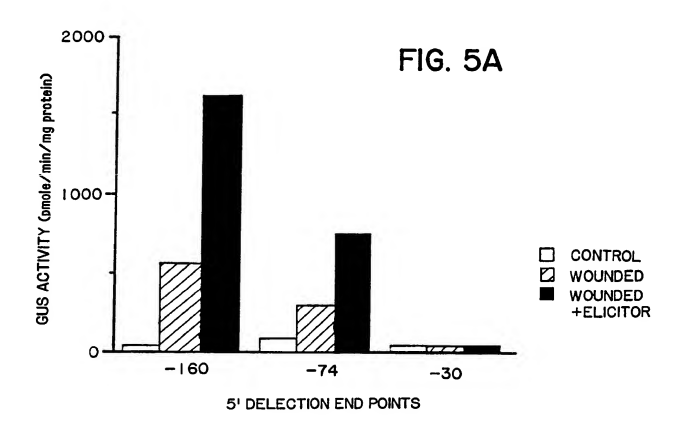


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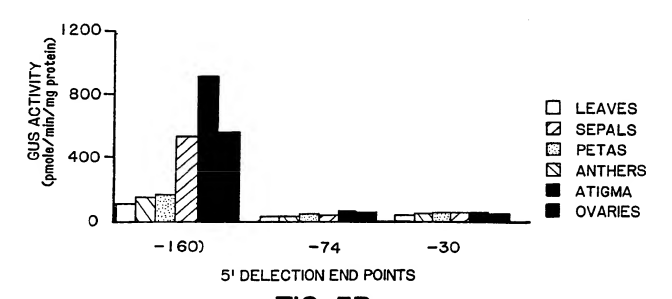


FIG. 5B

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INTERNATIONAL SEARCH REPORT

International application No.
PCT/US92/04282

A. CLASSIFICATION OF SUBJECT MATTER IPC(5) :C12N 15/82, 15/56, 5/14, 9/24; C12Q 1/68; A01H 5/00; C07K 13/00 US CL :536/27; 435/320.1, 44, 69.1, 200, 6; 800/205; 530/350 According to International Patent Classification (IPC) or to both national classification and IPC									
	LDS SEARCHED								
Minimum d	ocumentation searched (classification system followe	d by classification symbols)							
	536/27; 435/320.1, 44, 69.1, 200, 6; 800/205; 530/3								
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched									
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) Please See Extra Sheet.									
C. DOCUMENTS CONSIDERED TO BE RELEVANT									
Category*	Citation of document, with indication, where a	ppropriate, of the relevant passages	Relevant to claim No.						
X,P	MGG, vol. 226, issued 21 June 1991, Zhu et al, "gene encoding a basic chitinase", pages 289-296, e		1-20						
X,P	Plant Science, vol. 76, issued 22 July 1991, Nishizawa et al, "Rice chitinase gene: cDNA cloning and stress-induced expression", p. 211-218, especially figure 2 and page 216.								
x —	Plant Molecular Biology, vol. 16, issued March 1991, Huang et al, "Nucleotide sequence of a rice genomic clone that encodes a class I endochitinase", p. 479-480, entire document.								
Y			8-19						
Y	8-14								
Y	Physiologia Plantarum, vol. 79, issued July 1990, Jacobsen et al, "Characterization of two antifungal endochitinases from barley grain", p. 554-562, entire document.								
X Furth	ner documents are listed in the continuation of Box C	See patent family annex.							
"A" do	ecial categories of cited documents: cument defining the general state of the art which is not considered be part of particular relevance	"T" Inter document published after the inte date and not in conflict with the applica principle or theory underlying the inve	tion but cited to understand the						
"E" can	rijer document published on or after the international filing data current which may throw doubts on priority claim(s) or which is	"X" document of particular relevance; the considered movel or cannot be consider when the document is taken alone							
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"P" do	name current published prior to the international filing date but later than	being obvious to a person skilled in the "&" document member of the same passet.							
	actual completion of the international search	Date of mailing of the international sea	rch report						
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Commission Box PCT	nailing address of the ISA/	Authorized officer MARY E. MOSHER, PH.D.	Money						
_	L. D.C. 20231	Telephone No. (703) 308-0196							

INTERN IONAL SEARCH REPORT

i. _ornational application No. PCT/US92/04282

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Electronic data bases consulted (Name of data base and where practicable terms used):

EMBL-NEW 5, Genbank 71, Genbank-NEW 5, UEMBL 30-71, N-GeneSeq 6, APS, Biosis. Search terms: promoter, expression, monocot, maize, barley, wheat, rice, lily, onion, au = Zhu Q, au = Lamb C, hevein, chitinase, plant, plants, gene, genes, sequence?, clon?; sequences corresponding to nucleotides 1836-1884, 1810-1884, 1724-1884 of seq. ID no. 1; sequence corresponding to sequence ID no. 3.

Form PCT/ISA/210 (extra sheet)(July 1992)*

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